CASE REPORT

Pulmonary Thromboembolism Complicating Acute Pancreatitis With Pancreatic Ascites: A Series of 4 cases

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ABSTRACT

Acute pancreatitis is an inflammatory disease often associated with local and systemic complications. Portosplenic and splanchnic vascular complications of acute pancreatitis are common, but extrasplanchnic vessel thrombosis is less commonly seen. Among them, pulmonary thromboembolism is a very rare complication to be encountered with. We report four cases of acute pulmonary thromboembolism in patients with acute pancreatitis superimposed on chronic pancreatitis. All the patients had abdominal pain on presentation and distention of abdomen during the course. Dyspnea was present in all the patients. All patients were found to have pancreatic ascites, whose association with pulmonary thromboembolism is reported only in two patients till date up to our knowledge. Two of them had deep vein thrombosis and rest two had no venous thrombosis. All of them were managed conservatively using subcutaneous heparin, intravenous fluids and analgesics. We provide the causative mechanism for occurrence of pulmonary thromboembolism in acute on chronic pancreatitis. We have also hypothesized pancreatic ascites as the possible cause for pulmonary thromboembolism and provide explanation for it. We conclude that pulmonary thromboembolism in acute pancreatitis has good prognosis if diagnosed timely. Whenever patient with pancreatic ascites presents with dyspnea, pulmonary thromboembolism must be ruled out.

INTRODUCTION

Vascular complications of pancreatitis are among the causes of morbidity and mortality, common being haemorrhage into a pseudocyst, erosion of the arteries, rupture of pseudocyst, venous thrombosis and formation of varices [1]. Acute pancreatitis can result into systemic inflammatory response which may lead to a number of vascular and pulmonary complications. Portosplenic and splanchnic venous thrombosis are very well described in acute pancreatitis. However extrasplanchnic involvement is less common [2], particularly pulmonary thromboembolism, being a very rarely reported complication of acute pancreatitis [3-6].

Here we report four cases of pulmonary thromboembolism in acute pancreatitis on the top of chronic pancreatitis. We found that all the patients had pancreatic ascites, its association with pulmonary embolism has been reported only in two patients in the literature up to our knowledge [3].

CASE SERIES

Case #1

A forty-six-year-old male presented with a history of moderate epigastric pain since 1 month, increased in severity since last 10 days. He developed distention of abdomen after 1 week but without jaundice and pedal edema. He started developing dyspnea on exertion and coughing in last 15 days without any chest pain. Patient was alcoholic, consumed country alcohol around 120 grams per day since 20 years and chronic tobacco chewer. Heart rate was 88/min, Blood pressure was 130/80 mmHg and no hypoxia on arterial blood gas analysis. Abdominal examination showed epigastric tenderness with presence of ascites. Liver function tests were normal with amylase of 356 IU/L (15-100 IU/L) and lipase of 560 IU/L (0-60 IU/L). On diagnostic paracentesis, there were 1100 white cells/mm$^3$ (80% lymphocytes), 4.3 g/dL proteins and serum albumin ascites gradient (SAAG) of 0.6. Ascitic fluid amylase was 5401 IU/L. Contrast Enhanced Computed Tomography (CECT) scan of abdomen was showing atrophic pancreas with parenchymal calcification, peripancreatic fat stranding, 3 mm pancreatic duct and multiple intraductal calculi along with ascites (Figure 1). Two Dimensional Echocardiogram (2D Echo) and Electrocardiogram (ECG) were normal. Acute thrombosis of lobar and segmental branches of bilateral pulmonary arteries was noted in CT angiography with wedge shaped bands in both lung bases consistent with lung infarcts (Figure 2). His lower limb and abdominal vessel doppler were not suggestive of any thrombus.

Subcutaneous heparin was started along with analgesics and fluids. Pancreatographic demonstrated a
leak in the neck of pancreas and pancreatic duct stenting was done for the same. Ascites improved after few days. Patient shifted on oral anticoagulation and discharged at day 15.

Case #2

A thirty-year-old male, chronic alcoholic admitted to emergency department with complains of severe epigastric pain since 1 month, which was continuous, boring, radiating to back. He developed generalised abdominal distention after 5 days. He developed dyspnea on exertion before 1 week. He had similar type of severe abdominal pain before 1 year and 3 months, each time requiring hospital admission. He had multiple episodes of mild to moderate intermittent epigastric pain in 1 year, for which he used to take oral analgesics. He was alcoholic taking around 100 grams of country liquor daily since 8 years. Heart rate was 100/ minute and blood pressure was 120/70 mmHg at admission. Tachypnea was present with a rate of 25/minute but without hypoxia. Per abdomen was tender with severe ascites. Decreased air entry in both lower zones was noted on respiratory examination. Hemogram, liver enzymes and renal function tests were normal. Amylase was elevated (718 IU/L), so as lipase (400 IU/L). Diagnostic paracentesis was done, showing 135 cells, 85% neutrophils. Fluid proteins were 4.1 g/dL and amylase was 9290 IU/L. Atrophic pancreas with a leak in pancreatic duct in the body, peripancreatic edema and fat stranding were found on CECT abdomen along with ascitis. ECG had sinus tachycardia. Moderate pulmonary arterial hypertension (PAH) with dilated right atrium and ventricle were noted on 2D echo. CT pulmonary angiography (CTPA) suggested bilateral distal pulmonary artery thrombosis, extending into ascending, descending and segmental arteries (Figure 3). However, lower limb doppler was normal.

He was started on subcutaneous heparin and warfarin along with fluids and analgesics support. Patient improved after 15 days with conservative management.

Case #3

Thirty-eight-year-old male was hospitalised with 3 months history of severe epigastric pain, radiating to back and generalised abdominal distention and 15 days history of bilateral pedal edema. He also had dyspnea since 15 days, associated with left sided pleuritic chest pain. Patient consumed 80 grams of beer daily since last 5 years. Heart rate was 112/minute and blood pressure was 110/70 mmHg on admission. Abdomen was tender and distended with positive signs of free fluid and 5cm x 5cm bulge was palpated in epigastrium. Other systems were unremarkable. Serum amylase (1438 IU/L) and lipase (485 IU/L) were found elevated. Liver enzymes and hemogram were normal. Diagnostic ascitic tapping was done showing 3200 leucocytes / mm$^3$ (70% polymorphs), 3.6 g/dL of proteins and SAAG of 0.7. Fluid amylase was > 10,000 IU/L. CECT abdomen was suggestive of acute necrotising pancreatitis with a large pseudocyst with portal vein and superior mesenteric vein thrombosis.
Echocardiogram revealed moderate pulmonary arterial hypertension. Lower limb venous doppler suggested distal left saphenofemoral vein and popliteal vein thrombosis. To evaluate the cause for dyspnea with pleuritic chest pain, we ordered CT pulmonary angiography which showed left pulmonary artery thromboembolism with adjacent lung infarcts.

Patient was put on intravenous fluids, antibiotics and subcutaneous heparin. Symptoms decreased within a few days and patient was discharged on oral anticoagulation after 22 days.

Case #4

Thirty-five-year-old male presented with complain of severe epigastric pain since 15 days, radiating to back, followed after 3 days by abdominal distention and bilateral pedal edema, but more on left side. He had mild dyspnea since 4 days. He had an episode of acute pancreatitis with pleural effusion 3 months back. He used to drink 150 grams of country liquor since last 15 years. At admission, abdominal examination showed severe tenderness with ascites. Serum amylase was elevated (645 IU/L) with normal renal and liver function tests. Ascitic fluid examination revealed cells 250 cells/mm$^3$ (65% lymphocytes), 1.94 g/dL protein, SAAG of 0.6, amylase of 5286 IU/L and lipase >10,000 IU/L. CECT abdomen suggested acute on chronic pancreatitis with around 4 cm pseudocyst in body communicating with pancreatic duct with ascites (Figure 4). Left sided greater saphenous vein and left anterior tibial vein were found thrombosed on doppler. Despite of normal 2D echo and electrocardiography, CT pulmonary angiography showed bilateral segmental pulmonary artery thrombosis with right sided lung infarct (Figure 5).

Patient started on heparin, intravenous octreotide and fluids. Clinical improvement seen after 10 days and discharged at day 25.

DISCUSSION

Approximately one quarter of the patients with acute and chronic pancreatitis develop vascular complications [1]. Most common are haemorrhage into a pseudocyst, erosion of the arteries, rupture of pseudoaneurysm, venous thrombosis and formation of varices. Among them, venous thrombosis is a described complication in pancreatitis, particularly in acute pancreatitis [1].

The mechanism of venous thrombosis in acute pancreatitis is often multifactorial. First and the most frequently described mechanism is the inflammatory response associated with acute pancreatitis leading to a prothrombotic state, which makes vascular events more likely. Second, involvement of the vein by surrounding edema and cellular infiltration may cause venous thrombosis [7-9]. Third, compression of the vein by a pseudocyst or enlarged pancreatic parenchyma in turn precipitates vascular event [10]. Fourth, even a rupture of cyst into a vein also precipitates thrombosis [6]. At microvascular level, the mechanism of vascular injury may be due to intrinsic endothelial damage due to inflammatory mediators [11]. Involvement of veins in vascular thrombosis is variable. Portal vein, splenic vein and superior mesenteric vein are the most commonly involved vessels [12, 13]. Other vessels involvement have also been reported (Table 1) [15].

The pathogenesis of pulmonary embolism is also multimodal. It could be either due to thrombus detached from deep leg veins or sometimes inferior vena cavae [2]. Ohta et al. and Goenka et al. have reported an association of simultaneous inferior vena cavae thrombosis and pulmonary embolism with acute pancreatitis [6, 16]. Another case of acute pancreatitis with involvement of inferior vena cavae and pulmonary arteries in antiphospholipid antibody syndrome has been published [17]. Renal vein thrombus can also eventually cause pulmonary embolism. Only 2 cases have been reported so
Table 1. Thrombosis of different veins in acute pancreatitis.

<table>
<thead>
<tr>
<th>Author</th>
<th>Year</th>
<th>Patients</th>
<th>Involvement of veins</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peillon C [15]</td>
<td>1991</td>
<td>1</td>
<td>Inferior vena cava</td>
</tr>
<tr>
<td>Ohta H [6]</td>
<td>2002</td>
<td>1</td>
<td>Inferior vena cava</td>
</tr>
<tr>
<td>Ma SK [14]</td>
<td>2002</td>
<td>1</td>
<td>Inferior vena cava, renal vein</td>
</tr>
<tr>
<td>Parikh H [2]</td>
<td>2012</td>
<td>1</td>
<td>Superior and Inferior vena cava, bilateral Subclavian veins, Internal jugular and Iliac veins</td>
</tr>
<tr>
<td>Zhang Q [4]</td>
<td>2012</td>
<td>1</td>
<td>Pulmonary embolism</td>
</tr>
<tr>
<td>Phatak S [18]</td>
<td>2012</td>
<td>1</td>
<td>Inferior vena cava and pulmonary embolism</td>
</tr>
</tbody>
</table>

In an experimental study on pulmonary microvasculature in acute pancreatitis, mild pancreatitis showed abruptly terminating vessels. In more severe pancreatitis, there was substantial loss of vascular density, tortuosity, and abrupt terminations of those vessels present. This suggests that microvascular abnormalities are responsible for pulmonary infarction seen in acute pancreatitis [11].

Among four cases reported in present series, all the patients were having acute pancreatitis superimposed far [6, 14]. Inferior vena cavae and renal vein thrombosis results from either compression by psuedocyst or rupture of psuedocyst [6]. In recently published case series, pulmonary thromboembolism was proposed to be related to prothrombotic effect of inflammatory cytokines in acute pancreatitis [3]. Despite of vigorous efforts, no causative mechanisms have been elucidated in some cases [4].
on chronic pancreatitis. Etiology of chronic pancreatitis was alcohol in all of them. Predominant symptoms were abdominal pain (100%) and distention of abdomen (100%). All patients were symptomatic for pulmonary embolism, while in the series reported by Deiss et al. 2 out of 3 patients were found to have embolism incidentally [3]. Two patients (50%) had deep vein thrombosis, the presumed source for pulmonary embolism. One had left anterior tibial vein thrombosis while the other had left popliteal vein thrombosis. Rest of the patients did not have any deep vein or splanchnic vein thrombosis. However, causative vein thrombus is found only in nearly 50% cases of PTE. This is similar to our observation in the present series. Intrinsic endothelial damage may have played a role for vascular event in these cases.

Surprisingly, pancreatic ascites was a common association in all four cases while 2 out of 3 patients in the series reported by Deiss et al. also had pancreatic ascites [3]. Pancreatic ascites is one of the grave complications of pancreatitis, often along with pleural effusion, responsible for 20-30% mortality rates in some studies [18]. We hypothesize that pancreatic enzymes released in the ascitic fluid could have caused severe inflammation in the systemic veins. This may trigger cytokines to cause endothelial damage and venous thrombosis and thus, pulmonary embolism. This could be one of the possible explanation for association of pancreatic ascites with PTE. However, a direct causal relationship of pancreatic ascites with pulmonary embolism is a question of ongoing research.

In conclusion, pulmonary embolism is a rare association of acute and chronic pancreatitis. We should keep a high index of suspicion for pulmonary thoemboembolism in cases of acute on chronic pancreatitis with dyspnea, especially in the presence of pancreatic ascites which may confer a possible added risk for developing PTE.

Conflict of Interest

The authors declare that they have no competing interests.

References


